## PATENT COOPERATION TREATY

**PCT** 

REC'D 1 6 MAR 2006

# INTERNATIONAL PRELIMINARY REPORT ON PATENTABLETY

(Chapter II of the Patent Cooperation Treaty)

(PCT Article 36 and Rule 70)

Applicant's or agent's file reference	licant's or agent's file reference FOR FURTHER ACTION See Form PCT/IPEA/416				
PC-21016093					
THE PERSON NAMED IN COLUMN 1					
	T/SE2004/001644 10-11-2004 19-12-2003				
International Patent Classification (IPC) or national classification and IPC					
See Supplemental Box					
A 1:					
	Applicant				
CMS Contrast AB et al					
This report is the international pre Authority under Article 35 and tra	ansmitted to the applicant acco	ording to Article 2	s International Preliminary Examining 66.		
2. This REPORT consists of a total of		cluding this cover	sheet.		
3. This report is also accompanied by	y ANNEXES, comprising:				
		onu) a total of 3	sheets, as follows:		
	1 alaima and/or dra	wings which have	been amended and are the basis of this report		
and/or sheets	containing rectifications authors (rectifications)	orized by this Au	thornty (see Rule 70.10 and Section 607 61 and		
1 1 1 1 1 1		which this Author	ity considers contain an amendment that goes		
beyond the disconnection beyond the disconnect	isclosure in the international a	application as filed	d, as indicated in item 4 of Box No. I and the		
1		. 11	number of electronic carrier(s))		
b (sent to the Internation	onal Bureau only) a total of (1	ndicate type and i	number of electronic carrier(s)) and/or tables related thereto, in electronic		
form only, as indicate	, containing ted in the Supplemental Box R	d sequence fishing lelating to Sequen	ace Listing (see Section 802 of the		
Administrative Instr	uctions).				
4. This report contains indications r	relating to the following items	:			
Box No. I Basis	of the report				
Box No. II Priorit					
Box No. III Non-e	establishment of opinion with a	regard to novelty,	inventive step and industrial applicability		
	of unity of invention				
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Box No. VI Certai	in documents cited	iono supportung o			
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Box No. VIII Certai	in observations on the internat				
Date of submission of the demand	l i	Date of completio	n of this report		
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05-07-2005		06-03-200	6		
Name and mailing address of the IPEA		Authorized officer			
Patent- och registreringsverke	, 5.2				
BOX 5055 S-102 42 STOCKHOLM Malin Söderman/MP					
Facsimile No. +46 8 667 72 88		Telephone No. +46 8 782 25 00			

Form PCT/IPEA/409 (cover sheet) (April 2005)

International application No.

PCT/SE2004/001644

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Continuation of: Cover sheet

International patent classification (IPC)

A61K 49/06 (2006.01)

International application No.

PCT/SE2004/001644

Вох	No. I	Bas	is of the report	
1.	With 1	regard to t	he language, this report is based on:	
		the inter	national application in the language in which it was filed	
		o tronclo	tion of the international application into the language of a translation furnished for the purposes of:	,
			international search (Rules 12.3(a) and 23.1(b))	
		门	publication of the international application (Rule 12.4(a))	
			international preliminary examination (Rules 55.2(a) and/or 55.3(a))	
2.	furnis	shed to th are not an	o the <b>elements</b> of the international application, this report is based on (enterprise of the international application and invitation under Article 14 are referred nexed to this report):	replacement sheets which have been to in this report as "originally filed"
		the into	rnational application as originally filed/furnished	1
	$\boxtimes$	the des	cription:	and the flod formished
		pages	1-14	as originally filed/furnished
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		a seq	uence listing and/or any related table(s) – see Supplemental Box Relating to	Sequence Listing.
3	. [	The a	mendments have resulted in the cancellation of:	
			the description, pages	
			the claims, Nos.	
1			the drawings, sheets/figs	
-			the sequence listing (specify):	
			any table(s) related to the sequence listing (specify):	
	4.	This mad 70.2	report has been established as if (some of) the amendments annexed to the, since they have been considered to go beyond the disclosure as filed, as (c)).	nis report and listed below had not been indicated in the Supplemental Box (Rule
			the description, pages	
			the claims, Nos.	
1		Ī	the drawings, sheets/figs	
-		F	the sequence listing (specify):	
			any table(s) related to the sequence listing (specify):	
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International application No.
PCT/SE2004/001644

Box	No. II	Priority
1.	The lim	s report has been established as if no priority had been claimed due to the failure to furnish within the prescribed time it the requested:
		copy of the earlier application whose priority has been claimed (Rule 66.7(a)).
		translation of the earlier application whose priority has been claimed (Rule 66.7(b)).
2.	inv	is report has been established as if no priority had been claimed due to the fact that the priority claim has been found ralid (Rule 64.1). Thus for the purposes of this report, the international filing date indicated above is considered to be the evant date.
		al observations, if necessary:
	"Inci	priority is considered valid. Document Thomsen et al, eased Manganese Concentration in the Liver after Oral e", Academic Radiology, January 2004, vol. 11, no. 1, 38-44, is therefore of no relevance.
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International application No.

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Box No.	
The ques	stions whether the claimed invention appears to be novel, to involve an inventive step (to be non obvious), or to be industrially le have not been examined in respect of:
	the entire international application
$\boxtimes$	claims Nos. 21
becau	ise:
an	the said international application, or the said claims Nos. 21 relate to the following subject matter which does not require an international preliminary examination (specify): PCT Rule 67.1.(iv).: Methods for treatment of the human or imal body by surgery or therapy, as well as diagnostic thods.
	the description, claims or drawings (indicate particular elements below) or said claims Nosare so unclear that no meaningful opinion could be formed (specify ):
	the claims, or said claims Nos are so inadequately supported by the description that no meaningful opinion could be formed (specify ):
	no international search report has been established for said claims Nos.
	a meaningful opinion could not be formed without the sequence listing; the applicant did not, within the prescribed time
	furnish a sequence listing on paper complying with the standard provided for in Annex C of the Administrative Instructions, and such listing was not available to the International Preliminary Examining Authority in a form and manner acceptable to it. furnish a sequence listing in electronic form complying with the standard provided for in Annex C of the
	Administrative Instructions, and such listing was not available to the international Flemmary Examining Flemmary in a form and manner acceptable to it.
1	pay the required late furnishing fee for the furnishing of a sequence listing in response to an invitation under Rules 13ter, 1(a) or (b) and 13ter, 2.
	a meaningful opinion could not be formed without the tables related to the sequence listings; the applicant did not, within the prescribed time limit, furnish such tables in electronic form complying with the technical requirements provided for in Annex C-bis of the Administrative Instructions, and such tables were not available to the International Preliminary Examining Authority in a form and manner acceptable to it.
	the tables related to the nucleotide and/or amino acid sequence listing, if in electronic form only, do not comply with the technical requirements provided for in the Annex C-bis of the Administrative Instructions.
	See Supplemental Box for further details.

International application No.

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Box No. V Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

1. Statement

Novelty (N)	Claims Claims	1-20	YES NO
Inventive step (IS)	Claims Claims	1-20	YES NO
Industrial applicability (IA)	Claims Claims	1-20	YES NO

2. Citations and explanations (Rule 70.7)

Reference is made to the following documents:

D1: WO9811922 A2 D2: WO9702842 A1 D3: WO9605867 A2

D4: US4863898 A

D5: US6015545 A

The claimed invention relates to the use of a physiologically acceptable manganese (II) compound and an uptake promoter in the form of one or more amino acids for the manufacture of an MRI contrast composition for oral administration and MRI examination of the liver, in a ratio of Mn to promoter higher than that at which coordination compounds between Mn and promoter are formed to a substantial degree; an MRI contrast medium composition for such use; an MRI contrast medium kit; and a method for imaging of a mammalian liver using such contrast medium composition.

D1 describes an MRI contrast medium composition for use in a method for functional imaging of the gastrointestinal tract, see abstract. D1 also describes a method for rectal administration for obtaining images of the liver, see page 7, lines 5-18. In D1, manganese may be used in combination with a promoter, see page 8, line 14-page 9, line 25. The molar ratio of manganese to uptake promoter can be 1:0.2-1:50 or 1:1.5-1:5. The promoter can be, for example, alanine or aspartic acid.

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#### Supplemental Box

In case the space in any of the preceding boxes is not sufficient. Continuation of: Box  $\,\,V\,$ 

D2 describes a contrast medium that contains as active ingredient a manganese compound and an uptake promoter, see abstract. According to D2, 100 micromole/kg manganese(II)chloride tetrahydrate and 300 micromole/kg promoter can be used, see page 12.

D3 involves a contrast medium composition comprising a physiologically tolerable manganese compound and an uptake promoter and a physiologically tolerable carrier or excipient. The composition has a manganese concentration of 0.3 mM or is in a dosage unit form containing 300 micromole manganese, see abstract.

D4 relates to amino acid chelates having a ligand to divalent metal mole ratio of at least 2:1 for delivery to one or more specific tissue sites within a mammal, see abstract.

D5 describes a composition for use as a contrast medium being particularly suitable for imaging of the stomach, liver, bile duct and gall bladder, said composition comprising as an active ingredient a physiologically acceptable manganese compound and an uptake promoter, see abstract.

The cited documents represent the general state of the art. The invention defined in claims 1-20 is not disclosed by any of these documents.

The cited prior art does not give any indication that would lead a person skilled in the art to the claimed ratio of manganese to promoter. Therefore, the claimed invention is not obvious to a person skilled in the art.

Accordingly, the invention defined in claims 1-20 is novel and is considered to involve an inventive step. The invention is industrially applicable.

International application No.

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ox No. VI Certain documents cited			
Certain published documents (Rule 70	.10)		
Application No. Patent No.	Publication date (day/month/year)	Filing date (day/month/year)	Priority date (valid claim) (day/month/year)
Thomsen et al, "Incr after Oral Intake", no. 1, pages 38-44,	eased Mangane Academic Radi	se Concentratio	on in the Liver 2004, vol. 11,
Non-written disclosures (Rule 70.9)			Date of written disclosure
Kind of non-written disclo	Sure Date of nor (day)	n-written disclosure (month/year)	referring to non-written disclosur (day/month/year)

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#### CLAIMS

- 1. The use of a physiologically acceptable manganese (II) compound and an uptake promoter in the form of one or more amino acids for the manufacture of an MRI contrast composition for oral administration and MRI examination of the liver, in a ratio of Mn to promoter higher than that at which coordination compounds between Mn and promoter are formed to a substantial degree, wherein the molar ratio of Mn to promoter is in the range of from 2:3 to 3:1.
  - 2. The use according to claim 1, wherein said ratio is in the range of from 1:1 to 3:1.
  - 3. The use according to claim 2, wherein said ratio is in the range of from 2:1 to 3:1.

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- 4. The use according to any one of the preceding claims, wherein the dosage of manganese is in the range of from 25 to 150  $\mu$ mol/ kg body weight.
- 5. The use according to claim 4, wherein the dosage of manganese is in the range of from 50 to 125  $\mu$ mol/ kg body weight.
  - 6. The use according to claim 5, wherein the dosage of manganese is in the range of from 50 to 100  $\mu mol/\ kg$  body weight.
  - 7. The use according to any one of the preceding claims, wherein the uptake promoter is selected from the group consisting of alanine, valine, leucine, tryptophan, methionine, isoleucine, proline, phenylalanine, serine, glycine, threonine, cysteine, asparagine, glutamine, tyrosine, aspartic acid, glutamic acid, arginine, lycine and histidine.

2005-10-31 09:15 V:\\_NoOrganisation\CMC CONTRAST AB\PATENT\\_NoFacily\PCT\Z101cOb3\Z1016093 Amended claims TAN 20c5-10-31 1.doc

- 8. The use according to claim 7, wherein said promoter is selected from neutral amino acids including asparagine and aspartic acid.
- 9. The use according to claim 8, wherein said promoter is L-alanine.

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- 10. An MRI contrast medium composition for oral
  administration for examination of the liver comprising as
  an active ingredient a physiologically acceptable
  manganese (II) compound and an uptake promoter comprising
  one or more amino acids wherein Mn and the promoter are
  used in a molar ratio higher than that at which
  coordination compounds between Mn and promoter are formed
  to a substantial degree, wherein the molar ratio of Mn to
  promoter is in the range of from 2:3 to 3:1.
- 11. A composition according to claim 10, wherein 20 said ratio is in the range of from 1:1 to 3:1.
  - 12. A composition according to claim 11, wherein said ratio is in the range of from 2:1 to 3:1.
- 13. A composition according to any one of claims 10 to 12, wherein the dosage of manganese is in the range of from 25 to 150  $\mu$ mol/ kg body weight.
- 14. A composition according to claim 13, wherein the dosage of manganese is in the range of from 50 to 125  $\mu$ mol/ kg body weight.
- $15.\ A$  composition according to claim 14, wherein the dosage of manganese is in the range of from 50 to  $100\ \mu mol/\ kg$  body weight.

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- 16. A composition according to any one of claims 10 to 15, wherein the uptake promoter is selected from the group consisting of alanine, valine, leucine, tryptophan, methionine, isoleucine, proline, phenylalanine, serine, glycine, threonine, cysteine, asparagine, glutamine, tyrosine, aspartic acid, glutamic acid, arginine, lycine and histidine.
- 17. A composition according to claim 16, wherein said promoter is selected from neutral amino acids including asparagine and aspartic acid.

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- 18. A composition according to claim 17, wherein said promoter is L-alanine.
- 19. An MRI contrast medium kit comprising a first container accommodating a physiologically acceptable manganese (II) compound, and a second container

  20 accommodating an uptake promoter comprising one or more amino acids, and optionally, instructions for the use of the kit, the molar ratio of Mn to promoter being within the range of 2:3 to 3:1.
- 20. A kit according to claim 19, wherein said molar ratio, the dosage of manganese and/or said uptake promoter is (are) as defined in any one of claims 11 to 18.
- 21. A method for MRI of a mammalian liver using an MRI contrast medium composition according to any one of claims 10 to 18, said method comprising oral administration of an effective amount of said contrast medium composition.